

FORM PTO-1448 (REV. 7-200)	U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE	Atty. Docket No. 18896	Serial No. 10/535,434			
LIST OF PRIOR ART CITED BY APPLICANT (Use several sheets if necessary)		Applicants Kirby Siemering, et al.				
		Filing Date September 14, 2006	Group Art Unit 1634			
U. S. PATENT DOCUMENTS						
EXAMINER INITIAL*	DOCUMENT NUMBER	DATE	NAME	CLASS	SUBCLASS	FILING DATE (if appropriate)
U. S. PATENT PUBLICATION DOCUMENTS						
FOREIGN PATENT DOCUMENTS						
	DOCUMENT NUMBER	DATE	COUNTRY	CLASS	SUBCLASS	TRANSLATION
						YES NO
OTHER PRIOR ART (Including Author, Title, Date, Pertinent Pages, Etc.)						
/KS/	Van Hauwe P. et al., "Two Frequent Missense Mutations in Pendred Syndrome", <i>Human Molecular Genetics</i> , 7(7):1099-1104 (1998), XP-002454423					
↓	Leroy B.P. et al., "Spectrum of Mutations in <i>USH2A</i> in British Patients with Usher Syndrome Type II", <i>Experimental Eye Research</i> , 72(5):503-509 (2001), XP-002454423					
↓	Nájera C. et al., "Mutations in Myosin VIIA (<i>MYO7A</i>) and Usherin (<i>USH2A</i>) in Spanish Patients with Usher Syndrome Types I and II, Respectively", <i>Human Mutation</i> 20(1):1-7 (2002), XP-002454425					
↓	Bogazzi F. et al., "A Novel Mutation in the Pendrin Gene Associated with Pendred's Syndrome", <i>Clinical Endocrinology</i> , 52(3):279-285 (2000), XP-002454424					
↓	Weston M.D. et al., "Genomic Structure and Identification of Novel Mutations in Usherin, the Gene Responsible for Usher Syndrome Type Ila", <i>American Journal of Human Genetics</i> , 66(4):1199-1210 (2000), XP-002454426					
EXAMINER	/Katherine Salmon/		DATE CONSIDERED 06/29/2009			
* EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP 609; draw line through citation if not in conformance and not considered. Include copy of this form with next communication to applicant.						